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Spinning versus wobbling: how the brain solves a geometry problem

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Section Behavioral/Systems/Cognitive

Spinning versus wobbling: how the brain solves a geometry problem

Abbreviated title: Spinning versus wobbling

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Abstract

Oscillating an animal out-of-phase simultaneously about the roll and pitch axes ("wobble") changes continuously the head's orientation relative to gravity. For example, it may gradually change from nose-up, to ear-down, nose-down, ear-down and back to nose-up. Rotations about the longitudinal axis ("spin ") can change the head's orientation relative to gravity in the same way provided the axis is tilted from vertical. During both maneuvers the otolith organs in the inner ear detect the change in head orientation relative to gravity whereas the semicircular canals will only detect oscillations in velocity (wobble), but not any rotation at constant velocity (spin). Geometrically the whole motion can be computed based on information about head orientation relative to gravity and the wobble velocity. We subjected monkeys (*macaca mulatta*) to combinations of spin and wobble and found that the animals were always able to correctly estimate their spin velocity. Simulations of these results with an optimal Bayesian model of vestibular information processing suggest that the brain integrates gravity and velocity information based on a geometrically coherent three-dimensional representation of head-in-space motion.

Introduction

The perception of self-orientation and motion in space plays an important role in motor control of balance and locomotion. Over the last decade, evidence has accumulated that motor control involves central representations of “internal models” of movements (Ito 1989, Kawato 1999, Davidson and Wolpert 2005). Many modeling studies on spatial orientation and navigation also suggest that the brain maintains an internal estimate of three-dimensional motion in space, which is constantly updated to match afferent information from the vestibular, visual and proprioceptive sensory systems (Oman 1982, Droulez and Darlot 1989, Glasauer 1992, Merfeld et al. 1993, Merfeld 1995a, 1995b, Bos and Bles 2002, Zupan et al. 2002, MacNeilage et al. 2008, Green and Angelaki 2010). In a similar framework, Laurens and Droulez (2007, 2008), have recently proposed that the brain uses Bayesian inference to select the optimal estimate of the actual motion state.

Angular accelerations of the head are detected by the ampullary hair cells of the semicircular canals in the inner ear. However, these inertial sensors cannot detect prolonged rotations at constant velocity. If this rotation occurs about a tilted axis, the otolith organs do detect the body’s change in orientation relative to gravity (Off-Vertical Axis Rotation, i.e. OVAR, Fig. 1B). Primates and certain other species have developed mechanisms to efficiently use the vestibular afferent signals for gaze stabilization through vestibulo-ocular reflexes (VORs) (Guedry 1965, Young and Henn 1975, Raphan et al. 1981, Cohen et al. 1983, Harris 1987, Hess and Dieringer 1990; Hess and Angelaki 1993, Angelaki and Hess 1996a, b, Koshiro et al. 2002). Pertinent observations suggest that the brain estimates head motion in space by processing the vestibular information based on an internal model of head-in-space motion (see e.g. Merfeld et al 1999, Angelaki et al 2004). However, other authors have proposed that reflex eye movements could

simply result from appropriately filtering the otolith afferent signals (Hain 1986, Raphan and Schnabolk 1988, Raphan and Sturm 1991, Schnabolk and Raphan 1992, Raphan and Cohen 2002), without recourse to an internal model of motion.

To address this controversy, we have used a motion paradigm similar to one used earlier (Schor et al 1984) (cone motion, Fig. 1B), during which the head (and body) oscillates about the interaural and naso-occipital axes in phase quadrature such that it moves through the same sequence of orientations relative to gravity as during OVAR, yet without any rotation about the longitudinal axis. These oscillations activate the vertical semicircular canals that generate compensatory torsional and vertical eye movements (roll-pitch VOR) (Fig. 1C). Based on the synergy of vertical semicircular canal and otolith afferent activation, the internal model hypothesis predicts that no horizontal response will be generated during cone motion, in contrast to OVAR. A simple filtering of the otolith afferent signals would directly generate a horizontal VOR in both paradigms. We measured the three-dimensional (3D) VOR in rhesus monkeys during both OVAR and cone-motion, as well as in combinations of these two paradigms. The results of these experiments were simulated with a Bayesian model.

Methods

Figure 1 about here

Animal preparation and eye movement recording

Four female rhesus monkeys (*Macaca mulatta*) were chronically prepared with skull bolts for head restraint and dual search coils implanted under the conjunctiva for three-dimensional eye

movement recording as described in earlier studies (Hess 1990; Angelaki and Hess 1995). All procedures conformed to the National Institutes of Health Guide for the Care and Use of Laboratory Animals and were approved by the Veterinary Office of the Canton of Zurich.

Three-dimensional (3D) eye positions were measured using magnetic search coils (Robinson 1963) with an Eye Position Meter 3000 (Skalar, Delft, The Netherlands). Eye position was calibrated as described in Hess et al. (1992), digitized at a sampling rate of 833.33 Hz (Cambridge Electronic Device, Model 1401plus) and stored on a computer for off-line data analysis. Eye angular velocity vectors ($\boldsymbol{\Omega}$) were computed: the three components of $\boldsymbol{\Omega}$ represented the torsional (Ω_{tor}), vertical (Ω_{ver}) and horizontal (Ω_{hor}) angular eye velocity. Slow phase eye velocity was computed by detecting and removing quick phase movements using an algorithm similar to the one published by Holden (1992). In the following, the term “angular eye velocity” is synonymous to “slow phase angular eye velocity”.

Experimental set-up and stimulation protocols.

Monkeys were seated in a primate chair with the head restrained in upright position, so that the lateral semicircular canals were approximately earth-horizontal (stereotactic plane tilted about 15° nose-down). The primate chair was mounted on a computer controlled motorized 4-axis turntable, which was completely surrounded by a lightproof sphere of 1.6 m diameter. The inner wall of the sphere was covered with black dots of different sizes. The switching on and off of the inside illumination of the sphere was electronically controlled.

Spin and wobbling motion

In a first series of experiments, we compared the VOR elicited by a conventional OVAR paradigm (spin motion only, Fig. 1B, upper panel) with that due to a so-called cone motion paradigm (wobbling motion only, Fig. 1B, lower panel). The way spin and wobble motions superimpose in our paradigms can best be explained by the following line of thoughts. To generate the OVAR motion, the animal was rotated in a tilted position (tilt angle α) at constant velocity (ω_1) about its longitudinal axis (axis I in Fig. 1A, left). From the perspective of the animal, this motion changed its orientation relative to gravity or, equivalently, caused the gravity vector to rotate about its longitudinal axis (Fig. 1A, right). In order to induce wobble (Laurens et al. 2008) a further rotation was superimposed about an earth-vertical axis at a generally different constant velocity (ω_2 , space-fixed axis II in Fig. 1A, left). Since this second rotation was about an axis that was always aligned with gravity, it did not affect the animal's head orientation relative to gravity and therefore it did not modulate the otolith input. *In other words, the otolith stimulation was the same during all the experiments presented in this study.* From the animal's perspective, the angular velocity vector of this second motion (vector ω_2 , Fig. 1A, right) moved parallel to gravity, its tip describing a circle in the head yaw plane (Fig. 1A, right). The roll and pitch component of vector ω_2 therefore varied sinusoidally, creating a wobble motion. Note that the angular velocity ω_2 has a constant yaw (spin) component that adds or subtracts to the spin velocity ω_1 depending on the direction of rotation. Thus, counter-rotation about axis II will decrease the total spin velocity and increase the wobble velocity, eventually creating a pure wobble motion (cone), in which the head always faces the same direction in the horizontal plane. This can be visualized in Fig. 1B: the first axis of the platform rotates identically in the two paradigms, but the counter rotation of the platform during cone motion maintains the orientation of the head in the horizontal plane constant. Inversely, rotation about axis II in the same direction as about axis I will create a 'negative' wobble velocity and increase the spin velocity. The

resultant motion has a rotation component along the yaw axis, namely $S = \omega_1 - \omega_2 \cdot \cos \alpha$ (spin velocity), and a component $W = \omega_2 \cdot \sin \alpha$, (wobble velocity). Note that ω_1 , and α are constant. The spin (S) and the wobble (W) velocity are related to each other as follows:

$$S = \omega_1 - W / \tan \alpha \quad (\text{eq. 1})$$

Notice that during cone motion ($\omega_1 = \omega_2$), the head is always facing the same direction in the horizontal plane (Fig. 1B, lower panel) but the spin velocity of the head is in fact not zero because of the term $1 / \tan \alpha$ ($4^\circ/\text{s}$ for $\omega_1 = \omega_2 = 30^\circ/\text{s}$ with $\alpha = 30^\circ$).

OVAR motion was produced by tilting the axis I (Fig. 1A) by $\alpha = 30^\circ$ off vertical and rotating it at $\omega_1 = 30^\circ/\text{s}$ in darkness. Cone-motion was obtained by additionally counter-rotating the axis II at $\omega_2 = 30^\circ/\text{s}$. Combinations of these motions were obtained by rotating the axis II at velocities from $-60^\circ/\text{s}$ to $105^\circ/\text{s}$ in steps of $15^\circ/\text{s}$. In each animal, all combinations were tested in clockwise and counterclockwise direction (for which the signs of ω_1 and ω_2 were reversed). Motion duration was 90s for each trial. *Footnote: Animations of these protocols can be found at <http://www.vertigocenter.ch/laurens/SpinWobble.html>*

Spin stimulations during OVAR

In order to further investigate the interactions between spin and wobble motion in the VOR, we varied the sensory signals indicating a spin motion at the initiation of OVAR. This was done by applying stimuli that either activated the lateral semicircular canals or the horizontal optokinetic system. Initially, all animals were rotated in darkness at a velocity of $\omega' = (30 - \omega)^\circ/\text{s}$ around a vertical axis for 90s until the yaw VOR subsided. Then the rotation was rapidly changed to $30^\circ/\text{s}$, which elicited a yaw VOR with gain close to one. Two seconds after this acceleration step, the

animal was tilted towards 30° off-vertical (in 0.4 s, triangular velocity profile with $180^\circ/\text{s}^2$ acceleration) while it continued to rotate at $30^\circ/\text{s}$. In this manner, the induced sensory cues indicated a rotation in yaw (i.e. a spin motion) at any desired velocity ω when the OVAR was initiated, although the animal was actually spinning at $30^\circ/\text{s}$. Alternatively, in order to activate the visual system, the optokinetic sphere was rotated at $\omega' = (30 - \omega)^\circ/\text{s}$, while the animal rotated at $30^\circ/\text{s}$, both around a common vertical axis. During this phase preceding the OVAR, the animal thus rotated relative to the visual surround, i.e. the illuminated sphere at velocity ω . After 90s, the light was extinguished and the animal was tilted two seconds thereafter while continuing the rotation in yaw. The visual stimulation created a horizontal optokinetic nystagmus with a velocity close to ω , followed by an after response (optokinetic afternystagmus, OKAN) as soon as the light was extinguished. Both protocols were used to initiate OVAR trials with an initial yaw VOR/OKAN of $\omega = -60^\circ/\text{s}, -30^\circ/\text{s}, 0^\circ/\text{s}, 30^\circ/\text{s}, 60^\circ/\text{s}$. We also tested the VOR protocol at velocities $\omega = \pm 90^\circ/\text{s}$, but not the OKN/OKAN protocol, since these velocities exceeded the saturation velocity. Note that the total time between spin signal induction by vestibular or visual stimulation and completion of tilt was 2.4 s. This delay is considerably shorter than the time constant of the VOR and OKAN, thus the overlap of spin decay and OVAR onset was negligible. All animals underwent four trials at each of the 6 (7) velocities ω . All trials were initiated by tilting the animals through 30° from upright to either one out of the four orientations, i.e. nose-up, nose-down, left-ear down, and right-ear down. OVAR was maintained for 48s. Since the vestibular and optokinetic results were indistinguishable, we pooled the data for each of the four different head orientations.

Angular eye velocity analysis

The spin and wobble motions generated a constant-velocity horizontal VOR as well as vertical and torsional VOR velocities oscillating sinusoidally at a frequency of 1/12 Hz. This frequency corresponded to the rotation relative to gravity at $\omega_1 = 30^\circ/\text{s}$. In the following we call the constant horizontal response component spin or yaw VOR, and the oscillating vertical (Ω_{ver}) and torsional (Ω_{tor}) responses wobble VOR. The wobble VOR velocity was computed as follows: during wobbling at a velocity W , the head velocity in roll and pitch is $W \cdot (\cos(\theta), \sin(\theta))$. In this equation, θ is the orientation of the head relative to gravity, i.e. the component of gravity in the yaw plane of the head (nose-down: $\theta = 0$, right-ear down: $\theta = -90^\circ$, nose-up: $\theta = -180^\circ$, left-ear down: $\theta = -270^\circ$. Notice that θ decreases during clockwise rotation of the head relative to gravity, since this rotation causes gravity to rotate counterclockwise relative to the head). Thus for ideal compensation, the torsional and vertical VOR velocities should be $\Omega_{\text{tor}} = -W \cdot \cos\theta$ and $\Omega_{\text{ver}} = -W \cdot \sin\theta$. We computed the amplitude of the vector $(\Omega_{\text{tor}}, \Omega_{\text{ver}})$ as well as its phase relative to the ideal VOR, i.e. the velocity and the phase of the wobble VOR. Angular eye velocity during combinations of spin and wobble motion (Fig. 2C) was averaged over the last 48 s of stimulation. The angular eye velocity in Fig 3D. was averaged using a sliding window with a width of ± 0.5 s. We estimated the translational VOR during OVAR by an approach first used by Merfeld and colleagues (1999) (see also Fig. 5 and Results). According to this method, the yaw VOR during OVAR can be considered as a translational and a rotational VOR. Since gravity modulates sinusoidally along the interaural axis, the contribution of a possible translational VOR must be modulated in a sinusoidal manner as a function of head orientation, in contrast to the rotational VOR. We recorded the yaw VOR during OVAR trials, in which the animals were initially tilted towards nose-up, nose-down, left-ear down, and right-ear down. By averaging these trials, a contribution of the translational VOR is nullified, leaving a response representing the rotational VOR. Then, by subtracting the rotational VOR from the yaw VOR, we obtained the translational

VOR for each trial. By comparing trials at different head orientations, we estimated the influence of head orientation on the translational VOR at each point in time.

Bayesian modeling

The experimental results were compared to simulations performed with an optimal Bayesian model as described in (Laurens and Droulez 2007, 2008). This model computes an optimal estimate of head and body motion in space by assigning a probability to ‘every’ possible motion in space –in fact, the model only considers plausible motions in space, which are selected by a sampling process called particle filtering (Maskell and Gordon 2002). The probability depends on how well the sensory afferent signals that would be generated by a particular motion correspond to the actually received afferent signals, as well as to an a priori probability distribution of motions. The model assumes that the noise in the semicircular canals afferent signals limits the accuracy of the information provided about head angular velocity. In particular, the gain of the canals is zero during constant-velocity rotation, and therefore there is no information about steady-state yaw or spin velocity during OVAR and the cone-motion paradigm. In other words, all possible spin velocities are equally compatible with the afferent information of the semicircular canals. In contrast, the brain receives relatively accurate information about the wobble velocity, since wobble motion includes angular accelerations that efficiently activate the semicircular canals. Therefore the probability of a motion estimate that deviates from the wobble velocity sensed by the semicircular canals is low. The model also assumes that low translational accelerations are a priori more likely. Head motion relative to gravity and linear accelerations are sensed by the otolith organs, whose signals (F) are related to gravity (G) and linear head acceleration (A) according to $F = G - A$. The a priori probability on linear accelerations tends to favor motion estimates for which $A = G - F$ is small, i.e. for which the estimate of head

orientation relative to gravity (G) fully accounts for the afferent signals (F). In the process of Bayesian inference, each possible motion in space conforms to geometric constraints, and the orientation of the head in space (i.e. relative to gravity) is computed by integrating the estimated angular velocity over time. The prior on linear acceleration indirectly promotes rotation estimates that correspond to a rotation of the head relative to gravity as sensed by the otoliths. At the same time, the model assigns a high a priori probability to low angular velocities. This causes a small bias, which, however, plays a minor role in modeling the present paradigms. Finally, it is assumed that the visual-optokinetic pathways carry head velocity information relative to the environment with a certain signal-to-noise ratio that allows reproducing the phenomena of optokinetic nystagmus and afternystagmus (Laurens and Droulez 2007). We used the same model parameters as in (Laurens and Droulez 2008), i.e. $\sigma_{\Omega} = 40^{\circ}/s$, $\sigma_V = 10^{\circ}/s$, $\sigma_O = 7^{\circ}/s$, except for $\sigma_A = 1m/s^2$, which we adapted to the present experiments.

Results

Spin and wobble conventions

We have studied combinations of spin and wobble motion, which moved the head through the same orientations relative to gravity. The spin motion consisted of a constant-velocity rotation around the yaw axis (Fig. 1C), while the wobble motion consisted of oscillations around the pitch and roll axes (Fig. 1C). We measured the 3D VOR during these paradigms (Fig. 1C, lower panel). In order to be compensatory for a spin motion, the VOR should consist in constant-velocity horizontal eye movements ('spin VOR') whereas for wobble motion, it should exhibit oscillatory vertical and torsional eye movements ('wobble VOR').

Figure 2 about here

Angular eye velocity during the steady- state motion

In a first experiment, we tested whether the brain can correctly interpret the sensory signals during OVAR and cone motion. During OVAR, the head was rotating at a constant velocity of $30^\circ/\text{s}$ around the yaw axis (i.e. it was spinning). We found that monkeys exhibited a spin VOR (Fig. 2A), with an average velocity of $24 \pm 2.5^\circ/\text{s}$ (mean \pm SD between trials), close to the head velocity. A very small wobble VOR was also measured ($1.5 \pm 0.8^\circ/\text{s}$, $10 \pm 18^\circ$ phase lag). During the cone motion, the head was oscillating around the pitch and roll axes (wobble) with peak velocity of $15^\circ/\text{s}$ and also rotating in yaw (spinning) at a low velocity of $4^\circ/\text{s}$ (see eq. 1). We found a wobble VOR which closely matched the head wobble motion in phase and amplitude ($13.8 \pm 0.7^\circ/\text{s}$, $3 \pm 5^\circ$ phase lag), whereas the spin VOR was $2.7 \pm 0.5^\circ/\text{s}$ (wobble VOR, Fig 2B). This result was in line with our hypothesis that the spin VOR is based on an internal model of motion in space.

We tested this hypothesis further by submitting the animals to various combinations of spin and wobble motion, whose velocities are linked by eq. 1 (see Methods). Accordingly, each velocity combination represents a point on the line L in the spin-wobble plane (Fig. 2C). All of these combinations represent the same motion of the head relative to gravity. Therefore, the motion of the head relative to gravity as sensed by the otoliths during these experiments was geometrically consistent with any point located on line L. Finally, the semicircular canals directly detected the wobble component of the motion, while the brain received no direct sensory information about the spin velocity. We found that the eye movements measured during these combinations closely

corresponded to the head velocity provided the spin velocity remained below $30^\circ/\text{s}$. This was true even at the level of individual trials (grey lines in Fig. 2C). Outside of this range, the angular velocity of the VOR deviated from head velocity, suggesting that the central motion integration pathways reached their saturation point (Angelaki et al. 2000). This corroborates earlier findings that the brain estimates head-in-space motion by using a geometrically correct model of head motion, in which the estimate of angular velocity matches head motion relative to gravity (see Angelaki and Hess, 1995).

Figure 3 about here

Transient responses to spin stimulation

According to our working hypothesis, the internal estimate of head rotation should match the head orientation relative to gravity as sensed by the otoliths. Geometrically, this implies that the velocities of the spin and wobble VOR should closely correspond to eq.1, represented by the line L in Fig.2C. In the first series of experiments, we studied the VOR during steady-state motion. In this condition, the lateral semi-circular canals do not provide any information about the spin velocity. This velocity could nevertheless be computed as a function of the wobble velocities detected by the vertical semicircular canals as indicated by eq. 1. In a second series of experiments, we tested the hypothesis that sensory stimuli indicating a spin should reciprocally influence the wobble VOR according to eq. 1. At the beginning of OVAR, we therefore applied transient stimuli indicating a spin motion at various velocities (see *Spin stimulations during OVAR* in Methods) to test the hypothesis whether the animals would develop an internal estimate of wobble motion in an attempt to match the motion estimate predicted by eq. 1 (see line L in Fig 2C).

A first version of these experiments consisted in initiating OVAR without the usual preceding step-like activation of the semi-circular canals. In this condition, the spin VOR took several seconds to develop (Fig. 3A). During this period, we observed a wobble VOR that decayed while the spin VOR built up (in line with earlier reports, Hess and Angelaki 1999). Interestingly, we could amplify this effect by adding a negative step in spin rotation immediately before the onset of OVAR (see Methods). The initially negative spin VOR quickly reversed and reached the same steady-state level as during a normal OVAR trial (Fig. 3B). Simultaneously, the initial wobble VOR was enhanced and subsequently decreased as the spin VOR redeveloped. These phenomena are summarized in Fig. 3D: Starting from an initial condition in which the spin VOR was $-30^\circ/\text{s}$ and the wobble VOR zero (circle on Fig. 3D), the rapidly developing wobble VOR did bring the eye movements in about 2.5 s close to the theoretically expected line L (grey square). Then both the spin and wobble VOR converged towards a steady-state (star). In this phase, we observed a clear linear relationship between the spin and wobble VOR, as predicted by eq. 1. Finally, we applied positive spin velocity steps in a series of trials to induce a strong positive spin VOR. In the first few seconds of these paradigms, the wobble VOR showed an inverse phase relationship compared to the previous conditions (Fig 3C). This reversed VOR can be described as a negative wobble VOR (Fig. 3D) that also brought the eye velocity close to the expected line L. Subsequently, the spin and wobble VOR converged along a straight line towards steady state (star). Note that a certain degree of habituation occurred during these experiments. When an OVAR trial was performed in the first experiment (Fig. 2A,C), the steady state VOR reached an average of $24^\circ/\text{s}$ spin VOR and $1.5^\circ/\text{s}$ wobble VOR. In this second series of experiment, the steady-state reached $20^\circ/\text{s}$ spin and $2.5^\circ/\text{s}$ wobble VOR when the initial spin VOR was close to the real motion of the head (i.e. $30^\circ/\text{s}$). In other conditions (e.g. Fig. 3A-C), the steady-state VOR reached an average of $16.5^\circ/\text{s}$ spin and $3.5^\circ/\text{s}$ wobble, indicating that processing large spin signals

induced an additional habituation. Although the habituation affected the steady-state to a certain extent the linear relation between spin and wobble VOR during OVAR was preserved.

The line formed by the spin and wobble VOR had a similar orientation as the line L described by eq. 1 and it intersected the ordinate at about the same point. This suggests that it is indeed the kinematics represented by eq. 1 that governs the brain's adjustment of wobble velocity estimation (see line L in Fig 3D). However, although the experimental trajectory approached the line L, its slope was steeper (see line L in Fig 3D). In other words, the estimated the wobble VOR was only half in amplitude as that predicted by eq.1. Since there was no real wobble velocity of the head in this particular case, the estimated wobble which developed in an attempt to match the estimated motion of the head with the otolith signal was in conflict with the absence of semi-circular canal signals. We will interpret this lower amplitude as a trade-off between minimizing two mismatches in the Discussion.

Figure 4 about here

Bayesian modeling

We used the Bayesian model (Laurens and Droulez 2007, 2008) to compute the statistically optimal estimate of motion during our experimental paradigms. During OVAR, the motion estimate obtained from the simulated OVAR evolved towards a steady-state, for which the optimal estimate was spin rotation (Fig. 4A). During cone motion, the simulation produced an estimate of oscillations in pitch and roll, i.e. a wobble motion (Fig 4B), closely corresponding to the experimental findings (Fig. 2A, B). We simulated various combinations of spin and wobble motion and computed the spin and wobble responses based on optimal motion estimation. We

found that these responses formed a line close to the line L predicted by eq. 1 (Fig. 4C), in good agreement with experimental results. Moreover, during the first few seconds of the simulated OVAR response, we observed a wobble response (Fig. 4A) as during the experiments (Fig. 3A). Simulating OVAR trials with various initial spin amplitudes (Fig. 4D), we found that the estimated wobble was modulated as a function of initial spin velocity although it was lower than predicted by eq. 1, in agreement with the experimental findings. Altogether the simulations based on the Bayesian model could accurately reproduce a number of important aspects of our results.

Figure 5 about here

Linear acceleration estimation

So far, the interpretation of our results was based on the hypothesis that head motion relative to gravity is faithfully reported by the otolith organs. Head motion relative to gravity was the same in all our protocols, and corresponded to combinations of spin and wobble motion according to eq. 1 (see line L in Figs. 2C, 3D, 4C-D). Thus, we considered that the otolith information was in fact sufficient for the brain to know that the motion of the head corresponded to a motion according to line L. However, the otoliths are in fact sensitive to both the gravity vector (G) and linear acceleration (A) experienced during head translations, according to the equation $F = G - A$, in which F is the gravito-inertial acceleration which activates the otoliths. Therefore, although the animals were never translated in our experiment, the brain could have interpreted the variations of the otolith signals as the result of some incident linear accelerations of the head. A number of studies have shown that the brain, confronted to this ambiguity, tends to interpret the otolith signal as the consequence of gravity and minimize the linear acceleration estimate (Graybiel 1952, Guedry 1974, Paige and Seidman 1999). Therefore, our assumption that the brain attempts to develop a motion estimate that is coherent with head motion, as sensed by the otoliths, was a

justified simplification. However, the internal model hypothesis predicts that deviation of the angular motion estimate from the predicted line L (eq. 1) should result in a mismatch between the internal representation of gravity and the incoming otolith signals and lead to a non-zero estimate of translational head acceleration.

We tested this hypothesis by adopting a method used earlier by Merfeld and colleagues (1999). Linear acceleration along the interaural axis leads to a horizontal translational VOR that should superimpose to a rotational VOR with amplitude modulated by head orientation relative to gravity. At the frequency used in our experiment ($30^\circ/\text{s}$, i.e. 0.083 Hz motion of the head relative to gravity), this VOR should be in phase with the estimated acceleration of the head (Paige and Tokmo 1991). We illustrate this method by assuming that an animal is rotating in a counterclockwise direction. In an egocentric frame of reference, the gravity vector rotates clockwise around the animal's body vertical axis. If the estimate of gravity leads the actual head orientation (re gravity), then the translational VOR will take the form of rightward eye movements when the animals are in a nose-up position (Fig. 5A) and leftward movements in nose-down position (Fig. 5B). By summing up with the rightward spin VOR, the translational VOR should cause the total horizontal VOR to peak in nose-up orientation (Fig. 5A).

During the second experiment, we induced spin velocity signals at various velocities right before initiating the OVAR. In some conditions, these signals induced a spin VOR faster than $30^\circ/\text{s}$. We pooled the results whenever the spin VOR fell in between $40^\circ/\text{s}$ and $60^\circ/\text{s}$ (i.e. $50^\circ/\text{s}$ on average) and analyzed the modulation of the horizontal VOR as a function of head orientation (see Methods). We found that the VOR was faster under this condition in an orientation between nose-up and right-ear down (Fig. 5C, circle). This result was consistent with the hypothesis that the overestimation of the spin velocity caused the estimated gravity vector to lead the actual head

orientation, causing in turn a translational VOR. In a symmetric manner, the VOR peaked when the animals reached a nose-down orientation if the spin VOR was slower than $30^\circ/\text{s}$. Simulating the estimation of acceleration in these conditions with the Bayesian model, we found that the phase of the simulated estimate of interaural acceleration perfectly corresponded to the translational VOR measured in the experiments. In contrast, the average translational VOR during steady-state OVAR and cone motion remained close to zero (Fig. 5C, star).

Figure 6 about here

Principle of Bayesian estimation

The Bayesian model computes the probability distribution of three-dimensional motion over time. This high-dimensional process is difficult to visualize. However, it can be simplified by reformulating the inference in terms of spin and wobble motion. The OVAR stimulus consists only of spin motion about the z-axis (Fig. 6A) whereas the cone motion consists only of a wobble motion. In between these extremes there is a continuum of intermediate motion states (e.g. Fig 6B: mixture of $10^\circ/\text{s}$ wobble and $12.7^\circ/\text{s}$ spin motion). In this simplified version, we assume that the model ‘knows’ that the rotation is a combination of spin and wobble motion. Bayesian inference consists in computing the probability of all these combinations by combining three probability distributions. The first distribution is the a priori concerning the most likely distribution of angular velocities, which favors velocities close to zero (first column in Fig. 6A,B). The second is derived from the a priori about the distribution of translational accelerations, which favors low accelerations. Since motion estimates that contradict eq. 1 lead to an estimation of translational acceleration, this a priori favors motion estimates which are close to line L (second column). Finally, the third distribution comes from the semicircular canals, which can detect oscillations corresponding to the actual wobble motion with a certain uncertainty but

never rotations at constant velocity as the spin in our experiments, i.e. the uncertainty on spin velocity is infinite. Therefore the distribution of possible velocities according to the canals scatters around a vertical line W (third column, green line). The two lines L and W in Fig. 6 intersect at one point that corresponds to the real velocity of the head. The multiplication of the probability distributions according to the semicircular canal and otolith signals would lead to a final estimate centered at this intersection point. However, due to the uncertainty about these signals, the a priori knowledge influences the final estimate by shifting it somewhat towards zero, i.e. downward and rightward. This reduces the total velocity of the final estimate while maintaining it close to the line L. These examples illustrate how one can compute the steady-state velocity estimate using Bayesian inference. In addition, the time evolution of motion estimation during a spin-wobble paradigm can be obtained as follows: suppose for instance that the subject undergoes an OVAR motion ($30^\circ/\text{s}$ spin and no wobble), and consider a motion estimate corresponding to a certain position in the spin -wobble phase space (see open squares in Fig 6C). The effect of otolith signals will be to drive it towards the line L (no head translation, red arrows) while the effect of semicircular canal signals will be to attract it towards the ordinate axis (green arrows) since there is no wobble velocity during OVAR. Finally, the a priori information (not represented) will attract it towards the origin of the graph. Starting from a spin of $0^\circ/\text{s}$ (black square) or $60^\circ/\text{s}$ (grey square), the motion estimate quickly converges towards a line where the influence of the semicircular canal and otolith information is balanced and drive it slowly towards the final estimate (black circle). This process corresponds to the experimental measurements and to the simulations realized with the three-dimensional Bayesian model (Fig. 3D).

Discussion

We have demonstrated that the brain estimates head orientation in space in the cone-motion paradigm as well as in its variants with surprising accuracy. One rather suggestive way to explain this capacity is that the brain makes use of a geometrically appropriate three-dimensional internal model that represents motion in space. As shown in earlier studies, this type of reconstruction of motion in space is only feasible if the brain takes into account position and velocity information from both the otolith and the semi-circular canals (Hess and Angelaki 1993, Angelaki and Hess 1995, Angelaki et al 1999, Merfeld et al 1999, Green et al 2005). To push this analysis one step further we applied velocity steps of various amplitudes about the yaw axis while the animal was rotating in yaw at constant velocity in tilted position. These maneuvers induced conflicting sensory signals about the spin motion. To maintain geometrical consistency between the central rotation estimates and the head motion relative to gravity, we found that the animals developed an estimate of wobble motion, although they were in fact not wobbling. Yet, being in conflict with the missing modulation of vertical semi-circular canal activity the wobble estimate was lower than the amount necessary for perfect consistency with head motion relative to gravity. We used the Bayesian model to simulate the trade-off between geometrical consistency and sensory conflict, and found that the measured eye movements were close to the optimal solution. Finally, the internal model hypothesis predicted that the deviation from the estimated head motion relative to gravity and the actual gravito-inertial signals from the otolith organs should result in a non-zero estimate of translation. Indeed, the observed horizontal response can be interpreted as a translational VOR.

Our results support the notion that the brain integrates information about three-dimensional rotation velocity into a coherent estimate of head position in space. This estimate can be used to extract the inertial and gravitational components of the otolith afferent signals, in agreement with

previous studies (Merfeld et al. 1999, Angelaki et al. 1999, 2002, 2004). Our analysis suggests that conflicting motion signals are resolved by a process of optimal estimation, similar as shown in numerous previous studies (Ernst and Banks 2002, Weiss et al. 2002, MacNeilage et al. 2007, Angelaki et al. 2009, Fetsch et al. 2009) as well as in modeling work on vestibular information processing (Laurens et al. 2007, 2008). By demonstrating that the brain effectively uses geometrically consistent three-dimensional representations, this study supports the notion that the internal model hypothesis as formulated in the general context of motor control (Ito 1989, Kawato 1999, Davidson and Wolpert 2005) is also an important concept for understanding how the brain solves spatial orientation problems.

Previous work on the internal model hypothesis

Oman (1982) originally proposed that motion sickness arises when the brain fails to match an internal representation of motion with the sensory afferent inflow. This hypothesis inspired three-dimensional modeling such as the work of Droulez and Darlot (1989), which generates optimal motion estimates by using a gradient ascent procedure to minimize cost functions. The functions are designed in such a way that they favor estimates which are both geometrically consistent and match sensory inputs. Thus, the principles are the same as those underlying the Bayesian model. Later models by Merfeld (1993, 1995a,b), Glasauer (Glausauer and Merfeld 1997) and Zupan (Zupan et al. 2002) implemented the internal model in a computationally more efficient ad hoc architecture, which compares the expected sensory inputs with the effective afferent inflow. Any mismatch is used to correct the estimate of motion, which is computed according to an internal model of three-dimensional motion. Feedback loops ensure that the estimate is conforming to the sensory inflow (see also Bos and Bles 2002, MacNeilage and al 2008, Laurens and Angelaki

2011 for analyses and review of this framework). These loops can also generate a trade-off between different sources of sensory information when they are in conflict. All these models have been shown to successfully reproduce eye movements induced by off-vertical axis rotations.

Although the Bayesian model implements basically similar ideas as these previous models the crucial difference is that it performs the optimal estimation directly based on two simple assumptions: (1) the existence of an appropriate internal model of motion and (2) the existence of some measures quantifying or estimating the relative reliability of the various sensory inputs (e.g. based on the amount of noise in the afferent channels and possibly additional other factors). Thus, the critical advantage of using the Bayesian approach over previous models is that it allows one to test directly whether or not behavior can be explained by simple rules of optimal three-dimensional self-motion processing, as originally proposed by Oman (1982).

Other models of vestibular information processing

As an alternative to the internal model hypothesis, other authors (Hain 1986, Raphan and Sturm 1991, Angelaki 1992, Raphan and Cohen 2002) have proposed that the spin VOR measured during OVAR is produced by filtering otolith signals. However, these models predict that the otolith activation during cone motion, being identical to the activation during OVAR, should produce the same spin VOR, in contradiction with our results. Additionally, these studies focused on the spin response during OVAR, disregarding the wobble component that is pronounced at the onset of OVAR and cannot easily be accounted for by these models. In order to reproduce the present experimental results, these models would require substantial modifications, amongst which the addition of a specific mechanism which accurately cancels the spin VOR during cone

motion. Although this is of course possible, for instance by adapting the gravity-dependant velocity storage mechanism proposed by Raphan and Cohen (2002), it would make these models conceptually much more complex than the Bayesian approach, which is based on the internal model hypothesis. Furthermore, all the results of the experiments considered here agree perfectly well with the output of an optimal motion estimator. Therefore, adapting previous filtering models to these results would be equivalent to suggesting that the brain performs optimal estimation of head motion through a computationally and conceptually complex filtering process rather than by using an internal model.

Neural implementation

A variety of studies have addressed the issue of neural substrates that might be involved in constructing an internal model of self-motion in space. It has been demonstrated that the spin (yaw) VOR during OVAR is abolished after lesions of the nodulus and uvula (Angelaki1995), suggesting that the cerebellum plays a crucial role in building an internal model of self-orientation and motion. Such models seem to play an important role in many sensori-motor transformations (Ito 1989, Davidson and Wolpert 2005, Ito 2006, Green and Angelaki 2010) as well as possibly in Bayesian inference (Paulin 2005). Along these lines, Angelaki et al. (2004) have demonstrated that brain stem and cerebellar neurons integrate rotation information over time in order to extract the acceleration component from the otolith input. A realistic neural network architecture that integrates angular velocity in three dimensions was recently proposed by Green (Green et al. 2005).

Conclusion

Previous studies (Merfeld et al. 1999, Angelaki et al. 1999, 2004) showed that the brain uses angular motion information in order to disambiguate the otolith signals. Here we have created and tested a new family of motion paradigms, and demonstrated that the brain produces motion estimates which are geometrically consistent and can accurately be simulated by Bayesian inference.

Figure legends

Figure 1: Illustration of the OVAR and cone motion protocol. *A, left:* Schematic representation of the rotator used in the experiments. In all protocols, the subject was rotated about axis I at a constant velocity (ω_1). Simultaneous rotation about axis II generated a motion that is called wobble motion. Cone motion is generated when $\omega_1 = \omega_2$, as in B. *A, right:* Representation of (ω_1), (ω_2), and the gravity vector g in an egocentric frame of reference. *B:* Illustration of OVAR and cone motion paradigms. *upper panel:* In the OVAR paradigm, the head continuously changes its orientation relative to gravity while the subject rotates in tilted position at constant velocity about its longitudinal axis (LED = Left-Ear-Down, ND = Nose-Down, RED = Right-Ear-Down, NU = Nose-Up). *lower panel:* In the cone motion paradigm, the head changes its orientation relative to gravity in a similar way due to 90° out-of-phase pitch and roll oscillations. Thereby the subject faces always the same direction in the horizontal plane. Notice that the motion in both panels differs only by the counterrotation of the rotator. *C:* Representation of the three principal axes of head and eye movements. Curvilinear arrows indicate positive directions of head movements (head scheme) and eye movements (eye scheme).

Figure 2: Steady-state responses during OVAR, cone-motion and combinations of spin and wobble motion. *A, B:* Horizontal (blue), vertical (green) and torsional (red) eye velocity during three cycles of OVAR (“spin”) and cone motion (“wobble”). These panels show the response recorded during a representative trial. *C:* Responses to combinations of spin and wobble. *Open symbols:* head velocity; *solid symbols:* average horizontal angular velocity of yaw VOR; *grey lines:* individual trials. The line L corresponds to eq. 1.

Figure 3: Transient responses to spin pulse-injections before OVAR onset. *A*: Spin and wobble VOR during the first three cycles of OVAR initiated without any spin signal. *B*, *C*: Spin and wobble VOR during OVAR after injection of a negative (*B*) or positive (*C*) spin. *D*: Spin as a function of wobble showing an S-shaped behavior after an initial spin VOR of $-30^\circ/\text{s}$ (*blue circle and trace*) or after an initial spin VOR of $90^\circ/\text{s}$ (*diamond and magenta trace*). *Grey square*: VOR at $t = 2.5\text{s}$. *Blue star*: average steady-state VOR. For comparison, also line L is plotted (as in Fig. 2C). Traces in A-D are averages across all trials.

Figure 4: Bayesian model simulations. *A*: OVAR paradigm (compare with Fig. 2A and Fig. 3A). *B*: Cone-motion paradigm (compare with Fig. 2B). *C*: Simulated steady-state responses to combinations of spin and wobble motion (*blue line*) and comparison with the experimental results (as in Fig. 2C). *D*: Simulated transient responses to injection of spin pulses (as in Fig. 3D).

Figure 5: Translational VOR induced by in injection of spin stimuli before OVAR onset. *A*, *B*: Illustration of the acceleration estimate (a-estimate) generated when the estimated gravity (g-estimate) leads the actual gravity vector (g), during counter-clockwise rotation of the head, and of the superposition of the translational and angular (spin) VOR. *C*: Gain and phase versus head orientation of the translational VOR (blue, averaged across all trials) and interaural translational acceleration simulated with the Bayesian model (red), as a function of the spin VOR. The phase of the simulated translational acceleration is inverted in order to account for the fact that translational acceleration generates a VOR in the opposite direction.

Figure 6: Principles of Bayesian inference. The intensity plots represent the probability distributions according to the a priori, otolith and semicircular canal information in *A* for OVAR and in *B* for combinations of spin and wobble motion. The steady-state distribution (last column) is the product of the a priori distribution of angular velocity (first column) and the information

provided by the otoliths and the canals (2nd and 3rd column). The line L corresponds to eq. 1 as in the previous figures, the line W corresponds to the wobble velocity of the head. *Read diamonds*: real motion of the head; *Black circles*: the final estimate (last column). C: Evolution of the estimate over time during OVAR, starting from an initial value of zero (*black square, line*) or 60°/s spin (*grey square, line*). The arrows indicate the influence of canals and otoliths on the estimation at various points in time.

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